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(54) HEAT CURING-TYPE POWDER COATING COMPOSITION

(57)Abstract:

PROBLEM TO BE SOLVED: To provide a heat curing-type powder coating composition excellent in acid resistance and a finished outward appearance.

SOLUTION: The heat curing-type powder coating composition features comprising (I) a mixed resin between (A) an epoxy group-containing vinyl-based resin containing 20-70 wt.% of an epoxy group-containing vinyl-based monomer and having a glass transition temperature of 30°C to 100°C and (B) an epoxy group-containing vinyl-based resin containing 20-70 wt.% of an epoxy group- containing vinyl-based monomer and having a glass transition temperature of 30°C to 100°C wherein a solubility parameter of the resin (A) is smaller than that of the resin (B) and a difference between the solubility parameters of resins (A) and (B) is 0.5-1.7 and (II) one or more kinds of acid curing agents as curing resin component selected from the group consisting of a polycarboxylic acid curing agent, a carboxylic anhydride curing agent, and an anhydride curing agent from a polycarboxylic acid and a monocarboxylic acid as curing agents for the above resin basis.

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DETAILED DESCRIPTION

[Detailed Description of the Invention]

[0001]

[The technical field to which invention belongs] This invention is based on the thermosetting new powder-coatings constituent which can form the paint film excellent in weatherability, solvent resistance, acid resistance, and result appearance, and it is suitable for the fine-particles clear coating painted especially on a base coating.

[0002]

[Description of the Prior Art] finishing of a former and automobile shell plate -- the paint film which thermosetting powder coatings are painted as clear and formed there must be finished from the use, and must be filling the demand of appearance, solvent resistance, acid resistance, weatherability, etc. [0003] However, result appearance was good, the coexistence with the solvent resistance over low polar solvents, such as acid resistance, solvent resistance especially MINARARU spirits, and a xylol, was difficult, and the satisfactory thing was not obtained. [0004]

[Means for Solving the Problem] this invention person etc. came to complete a header and this invention for the ability of a heat-curing mold powder-coatings constituent with which result appearance is good with a constituent and may be compatible in acid resistance and solvent resistance to be offered, when mixed resin was used and a difference of the resin comrade's soluble parameter value used a thing of 0.5-1.7 as a base resinous principle for heat-curing mold fine particles as a result of inquiring wholeheartedly that the above-mentioned trouble should be canceled.

[0005] In 1 and heat-curing mold powder coatings, namely, a base resinous principle used with these powder coatings this invention Epoxy group content vinyl system resin within limits whose glass transition temperature which contains an epoxy group content vinyl system monomer in this resin 20 % of the weight to 70% of the weight as a radical polymerization nature partial saturation monomer component which constitutes following (resin I) (A) vinyl system resin is 30 degrees C - 100 degrees C (A), And it is mixed resin of epoxy group content vinyl system resin (B) of within the limits with a glass transition temperature of 30 degrees C - 100 degrees C which contains an epoxy group content vinyl system monomer in this resin 20 % of the weight to 70% of the weight as a radical polymerization nature partial saturation monomer component which constitutes (B) vinyl system resin. A soluble parameter of this resin (A) as a curing agent component of mixed resin whose difference of a soluble parameter of mutual resin smaller than a soluble parameter of resin (B) and is within the limits of 0.5-1.7, and the (II) above-mentioned base resin A heat-curing mold powder-coatings constituent characterized by containing one sort or two sorts or more of acid curing agents chosen from a multiplevalued carboxylic-acid curing agent, an anhydrous carboxylic-acid curing agent, and a curing agent of an anhydride of a multiple-valued carboxylic acid and a univalent carboxylic acid as a hardenability resingus principle. A soluble parameter value of epoxy group content vinyl system resin (A) A thing of 7.9-9.2 within the limits, [2 and a mixed resinous principle (I)] As a heat-curing mold powder-coatings constituent of a publication, 3, and epoxy group content vinyl system resin (A) to the above 1 whose

soluble parameter value of epoxy group content vinyl system resin (B) is mixed resin with a thing of 9.6-10.9 within the limits The above 1 which is 5 % of the weight - 60% of the weight of within the limits as a radical polymerization nature partial saturation monomer component from which a monomer of styrene and/or isobornyl (meta) acrylate constitutes this resin (A), or a heat-curing mold powder-coatings constituent given in 2, In 4 and a mixed resinous principle (I), it is involved in the above 1 whose mixed rate of the epoxy group content vinyl system (resin A) / epoxy group content vinyl system resin (B) is within the limits of 10 / 90 - 90/10 by weight ratio thru/or a heat-curing mold powder-coatings constituent given in any 1 term of 3.

[Embodiment of the Invention] In this description, a soluble parameter (SP) is the value calculated as follows.

[0007] A soluble parameter (SP) value is a value which it is measured by voice-sound-symbol titration and calculated according to the formula (Journal of Applied Polymer Science, 12, 2359, 1968) of following KW.SHU and J.M.CORBETT.

[0008]

$$SP = \frac{\int (VH) \delta H + \int (VD) \delta D}{\int (VH) \int (VD)}$$

(VH shows the volumetric fraction of a hexane among a formula, and, as for SP value of a hexane, and deltaD, in VD, the volumetric fraction of deionized water and deltaH show SP value of deionized water)

In the titration of a voice sound symbol, to the inside which dissolved 0.5g (solid content) of dried acrylic copolymers in acetone 10ml, in addition to n-hexane, the titration value [in / for titration value / in a voice sound symbol / H (ml) / reading and the voice sound symbol which adds deionized water into an acetone solution similarly] D (ml) is read, these are applied to the following formula, and VH, VD, deltaH, and deltaD are computed. In addition, SP values of each solvent are acetone:9.75, n-hexane:7.24, and deionized water:23.43.

[0009]

VH = H/(10 + H)

VD=D/(10+D)

deltaH=9.75x10/(10+H) +7.24xH/(10+H)

deltaD=9.75x10/(10+D) +7.24xD/(10+D)

Moreover, in this description, glass transition temperature is the value calculated by DSC (**** scanning calorimeter).

[0010] the glass transition temperature which contains [as a radical polymerization nature partial saturation monomer component from which the resinous principle (A) and resin (B) component which are used for this invention constitute vinyl system resin in each other] an epoxy group content vinyl system monomer preferably in each resin 25 % of the weight to 60% of the weight 20 % of the weight to 70% of the weight -- 30 degrees C - 100 degrees C -- desirable -- 35 degrees C - 80 degrees C and number average molecular weight 1000-20000 -- it is resin of 1500-15000 within the limits preferably. [0011] If the amount of epoxy group content vinyl system monomer components contained in a resinous principle (A) and resin (B) becomes less than 20% of the weight, respectively, hardenability will worsen, on the other hand, if the amount of epoxy group content vinyl system monomer components exceeds 70 % of the weight, respectively, it will be finished, and appearance worsens.

[0012] If the glass transition temperature of a resinous principle (A) and resin (B) becomes less than 30 degrees C, respectively, the blocking resistance of powder coatings will worsen, and on the other hand, if glass transition temperature exceeds 100 degrees C, respectively, the result appearance of a paint film will worsen.

[0013] Since the result appearance of a paint film will fall if particles start welding, blocking resistance

worsens and number average molecular weight exceeds 20000, respectively, the powder coatings with which the endurance of a paint film will fall and will be acquired if the number average molecular weight of a resinous principle (A) and resin (B) becomes less than 1000, respectively are not desirable. [0014] As the above-mentioned epoxy group content vinyl system monomer, glycidyl (meta) acrylate, the glycidyl allyl compound ether, 3, 4-epoxycyclohexyl (meta) acrylate, beta-methyl glycidyl (meta) acrylate, allyl glycidyl ether, etc. are mentioned, for example, these monomers -- one sort -- or two or more sorts can be used, combining. Glycidyl (meta) acrylate and beta-methyl glycidyl (meta) acrylate are especially suitable.

[0015] the resin (A) and resin (B) which are used by this invention -- the difference of the soluble parameter of mutual resin with the soluble parameter of this resin (A) smaller than the soluble parameter of resin (B) and -- 0.5-1.7 -- the thing of 0.7-1.5 within the limits is used preferably. If the difference of a soluble parameter becomes less than 0.5, the acid resistance of a paint film, solvent resistance, etc. will worsen, on the other hand, if the difference of a soluble parameter exceeds 1.7, the compatibility of resin

will worsen and the result appearance of a paint film will fall.

[0016] Moreover, it is desirable as resin (A) 7.9-9.2, and that a soluble parameter value uses the thing of the range of 8.1-9.0 especially. The effect excellent in paint film engine performance, such as acid resistance and solvent resistance (especially xylol-proof nature), is demonstrated by going into the range which the soluble parameter value described above.

[0017] On the other hand, it is desirable as resin (B) 9.6-10.9, and that a soluble parameter value uses the thing of the range of 9.8-10.7 especially. The effect excellent in paint film engine performance, such as acid resistance and solvent resistance (especially xylol-proof nature), is demonstrated by going into

the range which the soluble parameter value described above.

[0018] As for the mixed rate of resin (A) and resin (B), in mixed resin (I), 10 / 90 - 90/10 especially 20 / 80 - 80/20 are desirable on the basis of both sum total weight %. Since the solvent resistance of a paint film etc. will fall if the blending ratio of coal of resin (A) becomes less than 10 % of the weight, and the acid resistance of a paint film etc. will fall if the blending ratio of coal of resin (B) exceeds 90 % of the weight, the blending ratio of coal of resin (A), on the other hand, exceeds 90 % of the weight and the blending ratio of coal of resin (B) becomes less than 10% of the weight, it is not desirable.

[0019] As a method of adjusting so that it may go into the range of the soluble parameter value which described above resin (A) and resin (B), when the radical polymerization nature partial saturation monomer of the remaining following and remaining others which blended the above-mentioned epoxy group content vinyl system monomer 20 % of the weight to 70% of the weight is suitably chosen so that it may go into the range of the above-mentioned soluble parameter value, and a radical carries out a

polymerization reaction, it can adjust.

[0020] Moreover, as a radical polymerization nature partial saturation monomer of the ** used by resin (A), it is desirable to use styrene and/or isobornyl (meta) acrylate ten to 50% of the weight especially preferably five to 60% of the weight. Since the acid resistance of sufficient paint film will fall if less than 5% of the weight, and the result appearance of a paint film will fall on the other hand if it exceeds 60% of the weight, it is not desirable. As other radical polymerization nature partial saturation monomers For example, vinyl aromatic compounds, such as styrene, alpha-methyl styrene, vinyltoluene, and alpha-chloro styrene; Methyl (meta) acrylate, Ethyl (meta) acrylate, n-butyl (meta) acrylate, and i-butyl (meta) acrylate, t-butyl (meta) acrylate, 2-ethylhexyl (meta) acrylate, n-octyl (meta) acrylate, DESHIRU (meta) acrylate, Lauryl (meta) acrylate, stearyl (meta) acrylate, Alkyl ester or annular alkyl ester of carbon numbers 1-24 of acrylic acids, such as cyclohexyl (meta) acrylate, isobornyl (meta) acrylate, and tricyclo deca nil (meta) acrylate, or a methacrylic acid etc. is mentioned.

[0021] The curing agents (II) used by this invention are one sort or two sorts or more of curing agents chosen from a multiple-valued carboxylic-acid curing agent (a), an anhydrous carboxylic-acid curing agent (b), and the curing agent (c) of the anhydride of a multiple-valued carboxylic acid and a univalent

carboxylic acid.

[0022] Aliphatic series dibasic acid, such as a suberic acid, an azelaic acid, a sebacic acid, undecane diacid, dodecane diacid, tridecane diacid, tetradecane diacid, pentadecane diacid, and eicosane diacid, is

as specifically as a multiple-valued carboxylic acid (a) used suitably. Specifically as an anhydrous carboxylic-acid curing agent (b), the acid anhydride of aliphatic series dibasic acid, such as a suberic acid, an azelaic acid, a sebacic acid, undecane diacid, dodecane diacid, tridecane diacid, tetradecane diacid, pentadecane diacid, and eicosane diacid, can use it suitably.

[0023] Specifically as a curing agent (c) of the anhydride of a multiple-valued carboxylic acid and a univalent carboxylic acid, what was obtained by making anhydrous aliphatic series dibasic acid, such as a suberic acid, an azelaic acid, a sebacic acid, undecane diacid, dodecane diacid, tridecane diacid, tetradecane diacid, pentadecane diacid, and eicosane diacid, and monobasic acids, such as a palmitic

acid, stearin acid, and eicosanoic acid, can use it suitably.

[0024] In this invention powder coatings, the compound of others, such as a support inhibitor, a surface control agent, an anti-oxidant, an ultraviolet ray absorbent, UV stabilizer, an antiblocking agent, a floating regulator, an electrification control agent, a color pigment, a bulking agent, and a hardening accelerator, can be blended if needed in addition to the above-mentioned component.

[0025] this invention powder coatings can be manufactured by the well-known fine-particles manufacture method from the former. As the manufacture method of powder coatings, after carrying out the dryblend of mixed resin (I) and the curing agent (II) with dry-blending machines, such as a Henschel mixer, melting mixing of this blend object can be carried out with melting kneading machines, such as a biaxial melting kneading machine, and it can manufacture cooling, coarse grinding, pulverizing, and by carrying out screen filtration, for example. Moreover, reduced pressure (it heats if needed) desiccation of remaining as it is or the thing replaced by water can be carried out for the organic solvent solution which dissolves or distributes and becomes an organic solvent about mixed resin (I) and a curing agent (II) besides having described above, and an organic solvent and water can be removed, and it can solidify, and can manufacture coarse grinding, pulverizing, and by carrying out screen filtration. Furthermore, the organic solvent solution or aquosity liquid which dissolves or distributes and becomes an organic solvent about mixed resin (I) and a curing agent (II) besides having described above can be freeze-dried, and it can solidify, and can manufacture coarse grinding, pulverizing, and by carrying out screen filtration.

[0026] this invention powder coatings like the conventional powder coatings For example, steel, aluminum, On raw material front faces, such as a paint processing raw material which performed the primer, middle-coat paint, and under coat base paint of a surface treatment raw material or electropainting etc. which performed surface treatment, such as phosphoric acid zinc, to metal raw materials, such as zinc, tin, copper, and melting zinc galvanizing, and this metal raw material The thickness after printing usually carries out powder coating of the about 10-200 micrometers by the methods of application, such as electrostatic spray painting, frictional electrification spray painting, and fluidized bed coating, so that it may become the range of about 20-150 micrometers preferably. Subsequently For example, about 120 degrees C - 300 degrees C of hardening paint films can be preferably formed for about 10 - 60 minutes by baking for [for / about 20 minutes / -] 40 minutes preferably at the printing temperature of about 140 degrees C - 200 degrees C.

[Effect of the Invention] the resin (B) which uses the resin (A) component which uses this invention powder coatings in these coatings combining this resin -- since the soluble parameter is low -- a resin (A) component -- the upper layer of a hardening paint film -- and result appearance, such as the smooth nature on the front face of a paint film which a fine-particles particle carries out heat floating, and comes to form a paint film, and gloss, thinks that a good thing is obtained so that many resin (B) components in the lower layer may exist. Moreover, since it is a low polar paint film, the formed upper paint film cannot get wet easily to polar high corrosion matter, such as acid rain, and since it stops being able to permeate further easily into a paint film, it demonstrates the remarkable effect that paint film engine performance, such as acid resistance, is excellent. Moreover, since a high polarity paint film is formed in a lower layer, as the whole paint film, a low polar solvent, especially the bloating tendency to a xylol are controlled, and xylol-proof nature demonstrates as a result the remarkable effect of becoming good.

[Example] Hereafter, an example is lifted and this invention is explained to details. In addition, unless it refuses especially, weight criteria are expressed the "section" or "%."

[0029] Having taught the toluene 60 section to the reaction container equipped with the example thermometer of manufacture, the thermostat, the stirrer, the reflux condenser, and dropping equipment of resin (1), and blowing nitrogen gas into it, it heated at 105 degrees C and the mixed liquor of the styrene 20 section, the methyl methacrylate 10 section, the iso-butyl methacrylate 35 section, the glycidyl methacrylate 35 section, and the azobisisobutyronitril 4 section was dropped over about 3 hours. It was left at 105 degrees C after dropping termination for 1 hour, and further, the azobisisobutyronitril 0.5 section and the toluene 10 section were dropped over 1 hour, it was left at 105 degrees C after dropping termination for 1 hour, and the reaction was ended. Then, reduced pressure actuation was performed, the solvent in a system was removed, and resin (1) was manufactured. The soluble parameters of this resin (A-1) were 9.2, the glass transition temperature of 50 degrees C, and number average molecular weight 5000.

[0030] Having taught the toluene 60 section to the reaction container equipped with the example thermometer of manufacture, the thermostat, the stirrer, the reflux condenser, and dropping equipment of resin (2), and blowing nitrogen gas into it, it heated at 105 degrees C and the mixed liquor of the styrene 3 section, the methyl methacrylate 57 section, the n-butyl acrylate 5 section, the glycidyl methacrylate 35 section, and the azobisisobutyronitril 4 section was dropped over about 3 hours. It was left at 105 degrees C after dropping termination for 1 hour, and further, the azobisisobutyronitril 0.5 section and the toluene 10 section were dropped over 1 hour, it was left at 105 degrees C after dropping termination for 1 hour, and the reaction was ended. Then, reduced pressure actuation was performed, the solvent in a system was removed, and resin (2) was manufactured. The soluble parameters of this resin (2) were 10.3, the glass transition temperature of 55 degrees C, and number average molecular weight

[0031] Having taught the toluene 60 section to the reaction container equipped with the example thermometer of manufacture, the thermostat, the stirrer, the reflux condenser, and dropping equipment of resin (3), and blowing nitrogen gas into it, it heated at 105 degrees C and the mixed liquor of the styrene 20 section, the isobornyl methacrylate 25 section, the iso-butyl methacrylate 20 section, the glycidyl methacrylate 35 section, and the azobisisobutyronitril 4 section was dropped over about 3 hours. It was left at 105 degrees C after dropping termination for 1 hour, and further, the azobisisobutyronitril 0.5 section and the toluene 10 section were dropped over 1 hour, it was left at 105 degrees C after dropping termination for 1 hour, and the reaction was ended. Then, reduced pressure actuation was performed, the solvent in a system was removed, and resin (3) was manufactured. The soluble parameters of this resin (3) were 8.5, the glass transition temperature of 51 degrees C, and number average molecular weight 4800.

[0032] Having taught the toluene 60 section to the reaction container equipped with the example thermometer of manufacture, the thermostat, the stirrer, the reflux condenser, and dropping equipment of resin (4), and blowing nitrogen gas into it, it heated at 105 degrees C and the mixed liquor of the styrene 15 section, the methyl methacrylate 30 section, the iso-butyl methacrylate 20 section, the glycidyl methacrylate 35 section, and the azobisisobutyronitril 4 section was dropped over about 3 hours. It was left at 105 degrees C after dropping termination for 1 hour, and further, the azobisisobutyronitril 0.5 section and the toluene 10 section were dropped over 1 hour, it was left at 105 degrees C after dropping termination for 1 hour, and the reaction was ended. Then, reduced pressure actuation was performed, the solvent in a system was removed, and resin (4) was manufactured. The soluble parameters of this resin (4) were 9.5, the glass transition temperature of 52 degrees C, and number average molecular weight 5100.

[0033] Having taught the toluene 60 section to the reaction container equipped with the example thermometer of manufacture, the thermostat, the stirrer, the reflux condenser, and dropping equipment of resin (5), and blowing nitrogen gas into it, it heated at 105 degrees C and the mixed liquor of the styrene 11.5 section, the methyl methacrylate 33.5 section, the n-butyl acrylate 2.5 section, the iso-butyl methacrylate 17.5 section, the glycidyl methacrylate 35 section, and the azobisisobutyronitril 4 section

was dropped over about 3 hours. It was left at 105 degrees C after dropping termination for 1 hour, and further, the azobisisobutyronitril 0.5 section and the toluene 10 section were dropped over 1 hour, it was left at 105 degrees C after dropping termination for 1 hour, and the reaction was ended. Then, reduced pressure actuation was performed, the solvent in a system was removed, and resin (5) was manufactured. The soluble parameters of this resin (5) were 9.7, the glass transition temperature of 52 degrees C, and number average molecular weight 5000.

[0034] as a curing agent -- beer nova resin company make -- VXL-1381 (acid anhydride of dodecane

diacid) was used.

[0035] After carrying out the dryblend of the example hard resin of manufacture (1) 50 section of powder coatings (1), the hard resin (2) 50 section and the curing agent 40.0 section, and the benzoin 0.5 section with a Henschel mixer at a room temperature, melting kneading was carried out by the extruder. Next, after cooling, it pulverized by the pin disk, it filtered by 150 meshes, and powder coatings (1) obtained. Powder coatings (2) - (9) was performed like the example of manufacture of powder coatings (1) except changing combination, as shown in a table. A test result is shown in a table 1.

[A table 1] 表1

		,	実施側	翔鄉	翔網3	上類1	出数92	此較193	比較那4	上數第5	此較例6
粉体建料器号		1	. 2	3	4	5	6	7	8	9	
胎態(I)	觀	棚(A)	A-1	A-1	A-1	A-1	A-2	A-5	A-3	A-4	A-3
		棚(B)	A-2	A-2	A-2				A-2	A-2_	A-4
	皇	挪(A)	50	. 30	07	100	100	100	50	50 ·	50
	_	衛脂(B)	50	70	30				50	\$ 0	. 50
硬化剂(I)	VXL-		40	40	40	40	40	40	40	40	40
開 isp	樹脂	(A)	92	92	92	92	102	9.7	85	9.5	86
P80-7	間問		103	103	103		_	_	10.8	103	9.5
棚sp差	91 00	.,	1.1	1.1	1,1	1	-	_	1.8	QB	0.9
親親果	塗膜	外租	©	6	6	0	0	0	x	0	<u>@</u>
		11	. 0	. 0	0	0	×	Δ	0	X	0
,		口-16性	0	0	0	X	0	Δ	0	0	x

[0037] After having carried out air spray paint on the electrodeposted paint film which electrodeposited the epoxy system cation electrodeposition paint on the dull steel plate with a thickness of 0.8mm which performed preparation (1) coated-object zinc-phosphate chemical conversion of a sample so that it might become 20 microns of desiccation thickness, and was able to be burned so that it might become 25 microns of desiccation thickness about automobile middle-coat surfacer, and carrying out by the ability being burned for 30 minutes at 140 degrees C, wet sanding was carried out with the sandpaper of #400, ridge desiccation carried out, and it wiped with

[0038] (2) After having been burned so that it might become 20 microns of desiccation thickness, and carrying out automobile middle-coat surfacer on the electrodeposted paint film which electrodeposited the epoxy system cation electrodeposition paint on the dull steel plate with a thickness of 0.8mm which performed paint film creation condition zinc phosphate chemical conversion so that it might become 20 microns of desiccation thickness, and was able to be burned, wet sanding was carried out with the sandpaper of #400, and ridge desiccation was carried out. subsequently, serious -- cron base coat HM-22 (the Kansai Paint Co., Ltd. make, a metallic paint, trade name) was painted so that it might become about 15 microns by hardening thickness, printing hardening was carried out for 30 minutes at 140 degrees C with the oven, and it considered as the raw material for a trial.

[0039] Subsequently, powder coatings were electrostatically painted on the front face of this raw material so that thickness might become about 50 microns, and they were made it to carry out heat hardening at 160 degrees C with an oven for 30 minutes. The next trial was performed about the obtained color card. [0040] Paint-film appearance: The following criteria estimated the result appearance of a paint film from

a feeling of gloss, and a smooth feeling. O Although a good thing and O were inferior in smooth nature little, after heating for 15 minutes on the hot plate which trickled 0.4ml of sulfuric acids into the test color card acid-proof what [has a good feeling of gloss], thing [in which * is inferior a little], thing [in which x is inferior], and :40%, and was heated at 85 degrees C, it rinsed, the painted surface was observed and the following criteria estimated.

O: although it is completely changeless and normal to the thing **:painted surface, carry out the 0.5mL spot of the xylol on the xylene-proof [thing] nature:painted surface where few level differences are accepted in the boundary of the dropping section and the non-dropped section, and leave it at a room temperature for 30 minutes. Then, after wiping off a xylol by KAZE, visual assessment of the painted surface is carried out.

O: there is completely nothing with after.

**: A spot section outline can be checked.

x: Swelling clear to the spot section is accepted.

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CLAIMS

[Claim(s)]

[Claim 1] In heat-curing mold powder coatings, a base resinous principle used with these powder coatings Epoxy group content vinyl system resin within limits whose glass transition temperature which contains an epoxy group content vinyl system monomer in this resin 20 % of the weight to 70% of the weight as a radical polymerization nature partial saturation monomer component which constitutes following (resin I) (A) vinyl system resin is 30 degrees C - 100 degrees C (A), And it is mixed resin of epoxy group content vinyl system resin (B) of within the limits with a glass transition temperature of 30 degrees C - 100 degrees C which contains an epoxy group content vinyl system monomer in this resin 20 % of the weight to 70% of the weight as a radical polymerization nature partial saturation monomer component which constitutes (B) vinyl system resin. A soluble parameter of this resin (A) as a curing agent component of mixed resin whose difference of a soluble parameter of mutual resin smaller than a soluble parameter of resin (B) and is within the limits of 0.5-1.7, and the (II) above-mentioned base resin A heat-curing mold powder-coatings constituent characterized by containing one sort or two sorts or more of acid curing agents chosen from a multiple-valued carboxylic-acid curing agent, an anhydrous carboxylic-acid curing agent, and a curing agent of an anhydride of a multiple-valued carboxylic acid and a univalent carboxylic acid as a hardenability resinous principle.

[Claim 2] A heat-curing mold powder-coatings constituent according to claim 1 a thing of 7.9-9.2 within the limits and whose soluble parameter value of epoxy group content vinyl system resin (B) a soluble parameter value of epoxy group content vinyl system resin (A) is mixed resin with a thing of 9.6-10.9 within the limits for a mixed resinous principle (I).

[Claim 3] A heat-curing mold powder-coatings constituent according to claim 1 or 2 which is 5 % of the weight - 60% of the weight of within the limits as epoxy group content vinyl system resin (A) as a radical polymerization nature partial saturation monomer component from which a monomer of styrene and/or isobornyl (meta) acrylate constitutes this resin (A).

[Claim 4] A heat-curing mold powder-coatings constituent given in claim 1 whose mixed rate of epoxy group content vinyl system (resin A) / epoxy group content vinyl system resin (B) is within the limits of 10 / 90 - 90/10 by weight ratio in a mixed resinous principle (I) thru/or any 1 term of 3.

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(54) 【発明の名称】 熱硬化型粉体塗料組成物

(57)【要約】

(修正有)

【課題】 耐酸性、仕上がり外観に優れた熱硬化型粉体 塗料組成物を提供する。

【解決手段】 (I)(A)エポキシ基含有ビニル系単量体を20重量%~70重量%含有するガラス転移温度が30℃~100℃の範囲内のエポキシ基含有ビニル系 樹脂(A)、及び(B)エポキシ基含有ビニル系単量体を20重量%~70重量%含有するガラス転移温度30℃~100℃の範囲内のエポキシ基含有ビニル系 樹脂(B)の混合樹脂であって、該樹脂(A)の溶解性パラメーターが樹脂(B)の溶解性パラメーターよりも小さく、且つお互いの樹脂の溶解性パラメーターの差が0.5~1.7の範囲内である混合樹脂、及び(II)上記基体樹脂の硬化剤成分として、多価カルボン酸硬化剤、無水カルボン酸硬化剤、多価カルボン酸と1価カルボン酸の無水物の硬化剤から選ばれる1種もしくは2種以上の酸硬化剤を硬化性樹脂成分として含有することを特徴とする熱硬化型粉体塗料組成物。

【特許請求の範囲】

【請求項1】 熱硬化型粉体塗料において、該粉体塗料 で使用する基体樹脂成分が、下記樹脂(I)(A)ビニ ル系樹脂を構成するラジカル重合性不飽和単量体成分と して該樹脂中にエポキシ基含有ビニル系単量体を20重 量%~70重量%含有するガラス転移温度が30℃~1 00℃の範囲内のエポキシ基含有ビニル系樹脂(A)、 及び(B)ビニル系樹脂を構成するラジカル重合性不飽 和単量体成分として該樹脂中にエポキシ基含有ビニル系 単量体を20重量%~70重量%含有するガラス転移温 度30℃~100℃の範囲内のエポキシ基含有ビニル系 樹脂(B)の混合樹脂であって、該樹脂(A)の溶解性 パラメーターが樹脂(B)の溶解性パラメーターよりも 小さく、且つお互いの樹脂の溶解性パラメーターの差が 0.5~1.7の範囲内である混合樹脂、及び(II) 上記基体樹脂の硬化剤成分として、多価カルボン酸硬化 剤、無水カルボン酸硬化剤、多価カルボン酸と1価カル ボン酸の無水物の硬化剤から選ばれる1種もしくは2種 以上の酸硬化剤を硬化性樹脂成分として含有することを 特徴とする熱硬化型粉体塗料組成物。

【請求項2】 混合樹脂成分(I)が、エポキシ基含有 ビニル系樹脂(A)の溶解性パラメーター値が7.9~ 9.2の範囲内のものと、エポキシ基含有ビニル系樹脂 (B)の溶解性パラメーター値が9.6~10.9の範 囲内のものとの混合樹脂である請求項1に記載の熱硬化 型粉体塗料組成物。

【請求項3】 エポキシ基含有ビニル系樹脂(A)として、スチレン及び/又はイソボルニル(メタ)アクリレートの単量体が該樹脂(A)を構成するラジカル重合性不飽和単量体成分として5重量%~60重量%の範囲内である請求項1又は2に記載の熱硬化型粉体塗料組成物

【請求項4】 混合樹脂成分(I)において、エポキシ基含有ビニル系樹脂(A)/エポキシ基含有ビニル系樹脂(B)の混合割合が重量比率で10/90~90/10の範囲内である請求項1乃至3のいずれか1項に記載の熱硬化型粉体塗料組成物。

【発明の詳細な説明】

[0001]

【発明の属する技術分野】 本発明は、耐候性、耐溶剤性、耐酸性、仕上がり外観に優れた塗膜を形成できる新規な熱硬化性粉体塗料組成物によるものであり、特にベース塗料上に塗装される粉体クリヤー塗料に適したものである。

[0002]

【従来の技術及びその課題】 従来、自動車外板の上塗 りクリヤーとして熱硬化性粉体塗料が塗装され、そこに 形成される塗膜は、その用途から仕上がり外観、耐溶剤 性、耐酸性、耐候性等の要求を満たしていなければなら ない。 【0003】しかし、仕上がり外観が良好で、耐酸性と耐溶剤性特にミナラルスピリッツやキシロール等の低極性溶剤に対する耐溶剤性との両立は難しく、満足なものが得られていなかった。

[0004]

【課題を解決するための手段】 本発明者等は上記した問題点を解消すべく鋭意研究を行った結果、特に熱硬化型粉体用基体樹脂成分として、混合樹脂を使用し、その樹脂同志の溶解性パラメーター値の差が0.5~1.7のものを使用することにより、仕上がり外観が良好で、耐酸性と耐溶剤性を両立しうる熱硬化型粉体塗料組成物を提供できることを見出し、本発明を完成させるに至った

【0005】即ち、本発明は

- 1、熱硬化型粉体塗料において、該粉体塗料で使用する 基体樹脂成分が、下記樹脂
- (I)(A)ビニル系樹脂を構成するラジカル重合性不 飽和単量体成分として該樹脂中にエポキシ基含有ビニル 系単量体を20重量%~70重量%含有するガラス転移 温度が30℃~100℃の範囲内のエポキシ基含有ビニ ル系樹脂(A)、及び(B)ビニル系樹脂を構成するラ ジカル重合性不飽和単量体成分として該樹脂中にエポキ シ基含有ビニル系単量体を20重量%~70重量%含有 するガラス転移温度30℃~100℃の範囲内のエポキ シ基含有ビニル系樹脂 (B) の混合樹脂であって、該樹 脂(A)の溶解性パラメーターが樹脂(B)の溶解性パ ラメーターよりも小さく、且つお互いの樹脂の溶解性パ ラメーターの差が0.5~1.7の範囲内である混合樹 脂、及び(II)上記基体樹脂の硬化剤成分として、多 価カルボン酸硬化剤、無水カルボン酸硬化剤、多価カル ボン酸と1価カルボン酸の無水物の硬化剤から選ばれる 1種もしくは2種以上の酸硬化剤を硬化性樹脂成分とし て含有することを特徴とする熱硬化型粉体塗料組成物、 2、混合樹脂成分(I)が、エポキシ基含有ビニル系樹 脂(A)の溶解性パラメーター値が7.9~9.2の範 囲内のものと、エポキシ基含有ビニル系樹脂(B)の溶 解性パラメーター値が9.6~10.9の範囲内のもの との混合樹脂である上記1に記載の熱硬化型粉体塗料組 成物、
- 3、エポキシ基含有ビニル系樹脂(A)として、スチレン及び/又はイソボルニル(メタ)アクリレートの単量体が該樹脂(A)を構成するラジカル重合性不飽和単量体成分として5重量%~60重量%の範囲内である上記1又は2に記載の熱硬化型粉体塗料組成物、
- 4、混合樹脂成分(I)において、エポキシ基含有ビニル系樹脂(A)/エポキシ基含有ビニル系樹脂(B)の混合割合が重量比率で10/90~90/10の範囲内である上記1乃至3のいずれか1項に記載の熱硬化型粉体塗料組成物に係わる。

[0006]

【発明の実施の形態】 本明細書において、溶解性パラメーター(SP)は次のようにして求めた値である。 【0007】溶解性パラメーター(SP)値は、濁点滴 定によって測定されるものであって、下記K.W.SH U, J. M. CORBETTの式 (Journal of Applied Polymer Science, 12,2359,1968) に準じて計算した値である。

[0008]

 $\sqrt{(VH)} \delta H + \sqrt{(VD)} \delta D$

SP=

√ (VH) √ (VD)

(式中、VHはヘキサンの容積分率、VDは脱イオン水の容積分率、δHはヘキサンのSP値、δDは脱イオン水のSP値を示す)

濁点の滴定では、乾燥されたアクリル共重合体(固形分)0.5gをアセトン10mlに溶解した中に、n-ヘキサンに加えてゆき濁点での滴定量H(ml)を読み、同様にアセトン溶液中に脱イオン水を加えての濁点における滴定量D(ml)を読んで、これらを下記式に適用しVH、VD、δH、δDを算出するものである。尚、各溶剤のSP値はアセトン:9.75、n-ヘキサン:7.24、脱イオン水:23.43である。

[0009]

VH = H / (10 + H)

VD=D/(10+D)

 $\delta H = 9.75 \times 10 / (10 + H) + 7.24 \times H / (10 + H)$

 $\delta D = 9.75 \times 10/(10+D) + 7.24 \times D/$ (10+D)

また、本明細書において、ガラス転移温度は、DSC (示査走査熱量計)によって求めた値である。

【0010】本発明に用いられる樹脂成分(A)及び樹脂(B)成分はお互いにビニル系樹脂を構成するラジカル重合性不飽和単量体成分として夫々の樹脂中にエポキシ基含有ビニル系単量体を20重量%~70重量%、好ましくは25重量%~60重量%含有するガラス転移温度が30℃~100℃、好ましくは35℃~80℃、数平均分子量1000~20000、好ましくは1500~15000の範囲内の樹脂である。

【0011】樹脂成分(A)及び樹脂(B)中に含まれるエポキシ基含有ビニル系単量体成分量がそれぞれ20重量%未満になると硬化性が悪くなり、一方、エポキシ基含有ビニル系単量体成分量がそれぞれ70重量%を超えると仕上がり外観が悪くなる。

【0012】樹脂成分(A)及び樹脂(B)のガラス転移温度がそれぞれ30℃未満になると粉体塗料の耐ブロッキング性が悪くなり、一方、ガラス転移温度がそれぞれ100℃を超えると塗膜の仕上がり外観が悪くなる。【0013】樹脂成分(A)及び樹脂(B)の数平均分子量がそれぞれ1000未満になると塗膜の耐久性が低下し、また得られる粉体塗料は粒子同士が融着をおこし耐ブロッキング性が悪くなり、数平均分子量がそれぞれ20000を超えると塗膜の仕上がり外観が低下するので好ましくない。

【0014】上記エポキシ基含有ビニル系単量体としては、例えば、グリシジル(メタ)アクリレート、グリシジルアリルエーテル、3、4ーエポキシシクロヘキシル(メタ)アクリレート、βーメチルグリシジル(メタ)アクリレート、アリルグリシジルエーテル等が挙げられる。これらのモノマーは1種あるいは2種以上組み合わせて使用できる。特にグリシジル(メタ)アクリレート、βーメチルグリシジル(メタ)アクリレートが好適である。

【0015】本発明で使用する樹脂(A)及び樹脂

(B)は、該樹脂(A)の溶解性パラメーターが樹脂

(B)の溶解性パラメーターよりも小さく、且つお互いの樹脂の溶解性パラメーターの差が0.5~1.7、好ましくは0.7~1.5の範囲内のものが使用される。溶解性パラメーターの差が0.5未満になると、塗膜の耐酸性、耐溶剤性等が悪くなり、一方、溶解性パラメーターの差が1.7を上回ると樹脂の相溶性が悪くなり塗膜の仕上がり外観が低下する。

【0016】また、樹脂(A)として、溶解性パラメーター値が7.9~9.2、特に8.1~9.0の範囲のものを使用することが好ましい。溶解性パラメーター値が上記した範囲に入ることにより耐酸性、耐溶剤性(特に耐キンロール性)等の塗膜性能に優れた効果を発揮する。【0017】一方、樹脂(B)として、溶解性パラメーター値が9.6~10.9、特に9.8~10.7の範囲のものを使用することが好ましい。溶解性パラメーター値が上記した範囲に入ることにより耐酸性、耐溶剤性(特に耐キンロール性)等の塗膜性能に優れた効果を発揮する。

【0018】混合樹脂(I)において、樹脂(A)と樹脂(B)との混合割合は両者の合計重量%を基準として10/90~90/10、特に20/80~80/20が好ましい。樹脂(A)の配合割合が10重量%未満となり、そして樹脂(B)の配合割合が90重量%を越えると塗膜の耐酸性などが低下し、一方、樹脂(A)の配合割合が90重量%を超え、そして樹脂(B)の配合割合が10重量%未満になると塗膜の耐溶剤性などが低下するので好ましくない。

【0019】樹脂(A)及び樹脂(B)を上記した溶解性パラメーター値の範囲に入るように調整する方法としては、上記エポキシ基含有ビニル系単量体を20重量%~70重量%配合した残りの下記その他のラジカル重合性不飽和単量体を上記溶解性パラメーター値の範囲に入

るように適宜選択してラジカル共重合反応させることに より調整できる。

【0020】また、樹脂(A)で使用するその多のラジ カル重合性不飽和単量体としては、特にスチレン及び/ 又はイソボルニル (メタ) アクリレートを5~60重量 %、好ましくは10~50重量%使用することが好まし い。5重量%を下回ると十分な塗膜の耐酸性が低下し、 一方、60重量%を上回ると塗膜の仕上がり外観が低下 するので好ましくない。その他のラジカル重合性不飽和 単量体としては、例えば、スチレン、αーメチルスチレ ン、ビニルトルエン、αークロロスチレン等のビニル芳 香族化合物:メチル(メタ)アクリレート、エチル(メ タ) アクリレート、n-ブチル(メタ) アクリレート、 i-ブチル (メタ) アクリレート、 t-ブチル (メ タ) アクリレート、2ーエチルヘキシル(メタ) アクリ レート、nーオクチル(メタ)アクリレート、デシル (メタ) アクリレート、ラウリル (メタ) アクリレー ト、ステアリル (メタ) アクリレート、シクロヘキシル (メタ) アクリレート、イソボルニル (メタ) アクリレ ート、トリシクロデカニル (メタ) アクリレート等のア クリル酸またはメタクリル酸の炭素数1~24のアルキー ルエステルまたは環状アルキルエステル等が挙げられ

【0021】本発明で用いられる硬化剤(II)は、多価カルボン酸硬化剤(a)、無水カルボン酸硬化剤(b)、多価カルボン酸と1価カルボン酸の無水物の硬化剤(c)から選ばれる1種もしくは2種以上の硬化剤である。

【0022】多価カルボン酸(a)とては、具体的には、例えば、スベリン酸、アゼライン酸、セバシン酸、ウンデカン2酸、ドデカン2酸、トリデカン2酸、テトラデカン2酸、ペンタデカン2酸、エイコサン2酸等の脂肪族2塩基酸が好適に用いられる。無水カルボン酸硬化剤(b)としては、具体的には、例えば、スベリン酸、アゼライン酸、セバシン酸、ウンデカン2酸、ドデカン二酸、トリデカン2酸、テトラデカン2酸、ペンタデカン2酸、エイコサン2酸等の脂肪族2塩基酸類の酸無水物が好適に使用できる。

【0023】多価カルボン酸と1価カルボン酸の無水物の硬化剤(c)としては、具体的には、例えば、スベリン酸、アゼライン酸、セバシン酸、ウンデカン2酸、ドデカン二酸、トリデカン2酸、テトラデカン2酸、ペンタデカン2酸、エイコサン2酸等の脂肪族2塩基酸類とパルミチン酸、ステアリン酸、エイコサン酸等の1塩基酸類を無水化することで得られたものが好適に使用することができる。

【0024】本発明粉体塗料において、上記成分以外に 必要に応じてワキ防止剤、表面調整剤、酸化防止剤、紫 外線吸収剤、紫外線安定剤、ブロッキング防止剤、流動 調整剤、帯電制御剤、着色顔料、充てん剤、硬化促進剤 等のその他の配合物を配合することができる。

【0025】本発明粉体塗料は、従来から公知の粉体製 造方法により製造することができる。粉体塗料の製造方 法としては、例えば、混合樹脂(I)及び硬化剤(I I)をヘンシェルミキサー等の乾式混合機によりドライ ブレンドした後、このブレンド物を2軸溶融混錬機等の 溶融混練機により溶融混合し、冷却、粗粉砕、微粉砕、 篩沪過することにより製造することができる。また、上 記した以外に例えば、混合樹脂(I)及び硬化剤(I I)を有機溶剤に溶解もしくは分散してなる有機溶剤溶 液を、そのままもしくは水に置換したものを減圧(必要 に応じて加熱) 乾燥して有機溶剤や水を除去して固形化 し、粗粉砕、微粉砕、篩沪過することにより製造するこ とができる。更に、上記した以外に混合樹脂(I)及び 硬化剤(II)を有機溶剤に溶解もしくは分散してなる 有機溶剤溶液もしくは水性液を、凍結乾燥して固形化 し、粗粉砕、微粉砕、篩沪過することにより製造するこ とができる。

【0026】本発明粉体塗料は、従来の粉体塗料と同様に、例えば、鉄鋼、アルミニウム、亜鉛、錫、銅、溶融亜鉛メッキ等の金属素材、該金属素材にりん酸亜鉛等の表面処理を施した表面処理素材又は電着塗装などのプライマーや中塗り塗装や下塗りベース塗装を施した塗装処理素材などの素材表面に、焼付け後の膜厚が通常約10~200μm、好ましくは約20~150μmの範囲になるように、例えば、静電スプレー塗装、摩擦帯電スプレー塗装、流動浸漬塗装等の塗装方法で粉体塗装し、次いで、例えば、約120℃~300℃、好ましくは約140℃~200℃の焼付け温度で約10~60分間、好ましくは約20分間~40分間焼付けることにより硬化塗膜を形成することができる。

[0027]

【発明の効果】 本発明粉体塗料は、該塗料で使用する 樹脂(A)成分は該樹脂と組み合せて使用する樹脂

(B)よりも溶解性パラメーターが低いために、樹脂 (A)成分は硬化塗膜の上層にそして樹脂(B)成分はその下層に多く存在するように粉体粒子が熱流動して塗膜を形成するようになる塗膜表面の平滑性、光沢等の仕上がり外観が良いものが得られると考える。また、形成された上層塗膜は低極性塗膜であることから、酸性雨等の極性の高い腐食物質に対して濡れ難く、更に塗膜中に浸透し難くなるので耐酸性等の塗膜性能が優れるといった顕著な効果を発揮するものである。また、下層には高極性塗膜が形成されることから、塗膜全体としては低極性の溶剤、特にキシロールへの膨潤性が抑制され、結果として耐キシロール性が良好となるといった顕著な効果を発揮するものである。

[0028]

【実施例】 以下、実施例を揚げて本発明を詳細に説明 する。尚、特に断らない限り「部」または「%」は重量 基準を表す。

【0029】樹脂(1)の製造例

温度計、サーモスタット、撹拌器、還流冷却器および滴下装置を備えた反応容器に、トルエン60部を仕込み、窒素ガスを吹き込みながら、105℃に加熱して、スチレン20部、メチルメタクリレート10部、iso-ブチルメタクリレート35部、グリシジルメタクリレート35部、アゾビスイソブチロニトリル4部の混合液を約3時間かけて滴下した。滴下終了後105℃で1時間放置し、アゾビスイソブチロニトリル0.5部、トルエン10部を1時間かけて滴下し、滴下終了後1時間105℃で放置し、反応を終了した。その後、減圧操作を行い、系中の溶剤を除去し、樹脂(1)を製造した。該樹脂(A-1)は溶解性パラメーターが9.2、ガラス転移温度50℃、数平均分子量5000であった。

【0030】樹脂(2)の製造例

温度計、サーモスタット、撹拌器、還流冷却器および滴下装置を備えた反応容器に、トルエン60部を仕込み、窒素ガスを吹き込みながら、105℃に加熱して、スチレン3部、メチルメタクリレート57部、nーブチルアクリレート5部、グリシジルメタクリレート35部、アソビスイソブチロニトリル4部の混合液を約3時間かけて滴下した。滴下終了後105℃で1時間放置し、東に、アゾビスイソブチロニトリル0.5部、トルエン10部を1時間かけて滴下し、滴下終了後1時間105℃で放置し、反応を終了した。その後、減圧操作を行い、系中の溶剤を除去し、樹脂(2)を製造した。該樹脂(2)は溶解性パラメーターが10.3、ガラス転移温度55℃、数平均分子量5000であった。

【0031】樹脂(3)の製造例

温度計、サーモスタット、撹拌器、還流冷却器および滴下装置を備えた反応容器に、トルエン60部を仕込み、窒素ガスを吹き込みながら、105℃に加熱して、スチレン20部、イソボルニルメタクリレート25部、isの一ブチルメタクリレート20部、グリシジルメタクリレート35部、アゾビスイソブチロニトリル4部の混合液を約3時間かけて滴下した。滴下終了後105℃で1時間放置し、更に、アゾビスイソブチロニトリル0.5部、トルエン10部を1時間かけて滴下し、滴下終了した。その後、減日時間105℃で放置し、反応を終了した。その後、減日操作を行い、系中の溶剤を除去し、樹脂(3)を製造した。該樹脂(3)は溶解性パラメーターが8.5、ガラス転移温度51℃、数平均分子量4800であった。

【0032】樹脂(4)の製造例

温度計、サーモスタット、撹拌器、還流冷却器および滴下装置を備えた反応容器に、トルエン60部を仕込み、窒素ガスを吹き込みながら、105℃に加熱して、スチレン15部、メチルメタクリレート30部、isoーブチルメタクリレート20部、グリシジルメタクリレート35部、アゾビスイソブチロニトリル4部の混合液を約3時間かけて滴下した。滴下終了後105℃で1時間放置し、更に、アゾビスイソブチロニトリル0.5部、トルエン10部を1時間かけて滴下し、滴下終了後1時間105℃で放置し、反応を終了した。その後、減圧操作を行い、系中の溶剤を除去し、樹脂(4)を製造した。該樹脂(4)は溶解性パラメーターが9.5、ガラス転移温度52℃、数平均分子量5100であった。

【0033】樹脂(5)の製造例

温度計、サーモスタット、撹拌器、還流冷却器および滴下装置を備えた反応容器に、トルエン60部を仕込み、窒素ガスを吹き込みながら、105℃に加熱して、スチレン11.5部、メチルメタクリレート33.5部、nーブチルアクリレート2.5部、isoーブチルメタクリレート35 が 1 soーブチルメタクリレート35 が 1 soーブ・ボースイソブチロニトリル0.5部、トルエン10部を1時間かけて滴下し、滴下終了後1時間105℃で放置し、反応を終了した。その後、減圧操作を行い、系中の溶剤を除去し、樹脂(5)を製造した。該樹脂(5)は溶解性パラメーターが9.7、ガラス転移温度52℃、数平均分子量5000であった。

【0034】硬化剤として、ビアノバレジン社製VXL-13 81 (ドデカン2酸の酸無水物)を使用した。

【0035】粉体塗料(1)の製造例

固形樹脂(1)50部、固形樹脂(2)50部と硬化剤40.0部、ベンゾイン0.5部を室温でヘンシェルミキサーでドライブレンドした後、エクストルーダーで溶融混練した。次に冷却した後、ピンディスクで微粉砕し、150メッシュで沪過して粉体塗料(1)を得た。粉体塗料(2)~(9)は配合を表の様に変更する以外は粉体塗料(1)の製造例と同様に行った。試験結果を表1に示す。

[0036]

【表1】

表1

			親側	鄉	翔3	開	100002	城和3	战势4	雌鄉	城鄉6
粉体塗料醫号		1	2	8	4	5	6	7	8	9	
沿離(I)	馩	攤(A)	A-1	A-1	A-1	A-1	A-2	A-5	A-3	A∸4	A-3
		艦(8)	A-2	A-2	A-2				A-2	A-2	A-4
	量	挪(A)	- 50	30	70	100	100	100	50	50	50
		翻 (B)	\$ 0	70	. 30	,			8	50	· 50
副衛(I)	VXL-	1381	40	40	. 49	40	40	40	₽	40	40
開卸	樹脂	8	92	92	92	92	102	9.7	85	9.5	86
	間筒	8)	103	103	103	ļ	1	-	10.8	103	95
概能			1.1	1.1	1.1	Į	ı	_	1.8	0.8	Q .9
親親	塗膜	外観	8	0	0	0	0	0	X	9	· 6
	耐酸		0	0	0	0	X	Δ	0	Χ.	. 0
		10-14性	0	0	0	X	0	Δ	0	0	X

【0037】試料の調製

(1)被塗物

燐酸亜鉛化成処理を施した厚さ0.8mmのダル鋼鈑上にエポキシ系カチオン電着塗料を乾燥膜厚20ミクロンとなるように電着塗装し、焼き付けた電着塗膜上に自動車中塗りサーフェサーを乾燥膜厚25ミクロンとなるようにエアースプレー塗装し、140℃で30分間焼き付けした後#400のサンドペーパーで水研ぎし、水切り乾燥し、次いで石油ベンジンで拭いて脱脂し試験用被塗物とした。

【0038】(2)塗膜作成条件

燐酸亜鉛化成処理を施した厚さ0.8mmのダル鋼鈑上にエポキシ系カチオン電着塗料を乾燥膜厚20ミクロンとなるように電着塗装し、焼き付けた電着塗膜上に自動車中塗りサーフェサーを乾燥膜厚20ミクロンとなるように焼き付けした後#400のサンドペーパーで水研ぎし、水切り乾燥した。次いでマジクロンベースコートHM-22(関西ペイント株式会社製、メタリック塗料、商品名)を硬化膜厚で約15ミクロンとなるように塗装し、乾燥器で140℃で30分間焼付け硬化させ試験用の素材とした。

【0039】次いで該素材の表面に粉体塗料を膜厚が約50ミクロンとなるように静電塗装し、乾燥器で160℃で30分加熱硬化させた。得られた塗板について次の試験を行った。

【0040】塗膜外観:塗膜の仕上がり外観をツヤ感、 平滑感から次の基準で評価した。◎は良好なもの、○は 若干平滑性が劣るがツヤ感は良好なもの、△は若干劣る もの、×は劣るもの、

耐酸性:40%硫酸を試験塗板に0.4m1滴下し、85℃に加熱したホットプレート上で、15分間加熱した後、水洗し、塗面を観察し、次の基準で評価した。

○:全く変化のないもの

△: 塗面に異常はないが、滴下部と非滴下部の境界にわずかな段差が認められるもの

耐キシレン性:塗面上にキシロールを0.5mLスポットし、30分室温で放置する。その後、カーゼでキシロールを拭き取った後、塗面を目視評価する。

○:全く後付きがない。

△:スポット部輪郭が確認できる。

×:スポット部に明らかな膨潤が認められる。

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